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(57) Abstract

Method for detecting Campylobacter by PCR detection of DNA sequence, highly conserved between species lari, coli, jejuni and upsaliensis. Speciation between these four is possible as the PCR product is differentially cleaved by restriction endonucleases.

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- 1 -

DETECTION AND SPECIATION OF CAMPYLOBACTER

This invention relates to the detection and speciation of campylobacter bacteria, for example in clinical, environmental and food samples. In particular, this invention relates to a method of detecting whether a sample contains campylobacter and to a method of differentiating between the main campylobacter species jejuni, coli, upsaliensis and lari.

campylobacter species are recognised as the most frequent cause of bacterial gastroenteritis in the United Kingdom and many other countries throughout the world. In the U.K. approximately 90% and 10% of case isolates are identified as campylobacter jejuni and campylobacter coli respectively, plus a small number of other species such as campylobacter upsaliensis and lari. The majority of the infections are sporadic the source of which remains largely unknown although the importance of several vehicles is now recognised.

There is a known desire to be able to detect and differentiate species of campylobacter. However, it is also known that present campylobacter enrichment culture techniques lack sensitivity, making detection difficult. campylobacter jejuni does not multiply in foodstuffs and low numbers may be present together with a high background of indigenous microflora. Also, surface viable counts of campylobacter can decrease rapidly and cells that are potentially culturable are often lost before samples reach a laboratory for analysis. Another factor making detection problematic is that antibiotics used in culture enrichment media may damage already weakened campylobacter.

There are currently available assays for detection of a variety of food and water-borne pathogens; L.pneumophila,

- 2 -

V.vulnificus, enteroinvasive E. coli, Shigella; but no satisfactory method of detecting campylobacter or distinguishing between the four main campylobacter species is known.

A method of detecting campylobacter has been published by Giesendorf, B A J, et al in Applied and Environmental Microbiology, December 1992, pages 3804-3808. detects the species jejuni, Coli and lari, and produces similar results to conventional methods but in a reduced time. The method suffers from a number of drawbacks. It does not enable detection of the species upsaliensis. Further, the method employs polymerise chain reaction (PCR) techniques but nevertheless requires a short enrichment culture before the PCR can be employed. Further still, the primer used for the PCR does not have the precise homology with DNA sequences in the three campylobacter species that can be detected using the method.

Another method for detecting campylobacter jejuni and campylobacter coli is known from Wegmuller, B E et al, Applied Environmental Microbiology, vol. 59, part 7, 1993 pages 2161-2165. The described method detects only the species jejuni and coli.

In addition to the above-identified problems with detection and speciation of campylobacter, recent work on campylobacter jejuni suggests that in certain circumstances it enters a "non-culturable, viable form" when subjected to environmental stresses, such as pH or temperature extremes, increased oxygen tension or nutrient depletion. In this form, campylobacter infectivity is maintained but the organisms cannot be cultured. Thus there exists a need for the improvement of methods of detection of non-culturable forms of campylobacter. - 3 -

It is an object of this invention to provide a method of testing for the presence of campylobacter that enables more efficient detection and eliminates or mitigates the problems with existing techniques. It is a further object to provide a method of distinguishing the campylobacter species jejuni, coli, upsaliensis and lari.

Accordingly, in a first aspect the present invention provides a method of testing for the presence of campylobacter, e.g. in a clinical, environmental or food sample, comprising the steps of performing polymerise chain reaction (PCR) using primers adapted to amplify a region selected from (a) a sequence of at least 72 base pairs from sequence ID No. 1 and (b) a sequence having sufficient homology with (a) such that formation of PCR product is correlated with presence of campylobacter; and determining if any PCR product is formed.

It is preferred that sequence (b) has at least 75% homology with (a), preferably at least 90% homology and more preferably at least 95% homology. It is also preferred that the primers are at least 12 nucleotides in length, preferably between 19-22 nucleotides in length. In particularly preferred embodiments of the invention the primers consist of at least 12 contiguous nucleotides selected from (1) sequence ID NO:2 and sequence ID NO:3, (2) sequence ID NO:4 and sequence ID NO:5 and (3) sequences having sufficient homology with (1) or (2) such that formation of PCR product is correlated with presence of campylobacter

PCR has become a well known and established tool for DNA analysis. A single gene sequence can be marketed from a large amount of other DNA and amplified to provide a suitable quantity for analysis. The basis of todays PCR was first published in 1971 be Kleppe, E et al, J. Mol. Biol., 1971, 56, 341. Further significant details and improvements on the PCR

- 4 **-**

method have been added by Saiki, R K et al, science, 1985, 230, 1350 and Mullis, K B, sci. am. 1990, 262, 36.

As will be appreciated by a person of skill in the art familiar with the PCR, it is important to operate at a temperature suitable to ensure that the primers used are specific for the sequence desired to be identified and amplified. To this end it is convenient to carry out the PCR reaction using the method of the invention at temperatures of at least 40°C preferably at least 45°C and in a particularly preferred embodiment at 48-52°C.

The 1.9 kilobase fragment identified in sequence ID 1 is an underlying feature of this invention and has been found to be highly conserved between campylobacter isolates. The method of the invention confers the advantage that PCR product will only be detected when a campylobacter strain is found in the sample tested. The method also confers the advantage that it will detect non-culturable viable forms of campylobacter as well as viable cells. Thus the method is effective where other methods have not been able to detect any campylobacter.

It is preferable to use a primer sequence that will only bind to one specific region of sequence ID NO:1 and which will not engage in formation of primer dimers and thus contaminate the PCR. Examples of preferred primers for use in the method of the invention are shown in the sequence ID numbers 2 and 3 and sequence ID numbers 4 and 5. These primers form further aspects of the invention.

In a second aspect the invention provides a method of distinguishing between campylobacter species jejuni, coli, upsaliensis and lari in a DNA containing sample by performing PCR utilizing primers capable of amplifying a selected campylobacter DNA sequence, said sequence having restriction

endonuclease sites specifically associated with different campylobacter species and then testing for digestion of the PCR product by the specific restriction endonucleases.

Thus, campylobacter DNA that is differentially cleared by restriction endonucleases is amplified, subject to digestion by the endonucleases and identified as from a particular species.

Sequence ID NO:1 was isolated from Campylobacter jejuni and is known to have a particular characteristic pattern of cleavage by restriction endonucleases. Campylobacter coli, upsaliensis and lari contain sequences corresponding to sequence ID NO:1 that have altered patterns of cleavage characteristic of each species.

In an embodiment of the second aspect there is provided a method of distinguishing between campylobacter species jejuni, coli, upsaliensis and lari, e.g. in a clinical, environmental or food sample containing campylobacter, comprising the steps of:-

performing polymerise chain reaction (PCR) on the sample using primers adapted to amplify a region of DNA sequence ID No.1 that includes nucleotides 124-196, or using primers adapted to amplify a DNA region corresponding thereto; and

testing the PCR product for digestion by restriction endonucleases Alu I, Dra I and Dde I.

The method of the second aspect is advantageous because it enables accurate speciation between the four clinically most significant species. In particular, when amplifying region 124-196 of SEQ DI NO:1, the PCR product from campylobacter

jejuni is cleaved by all three restrictions endonucleases, whereas the PCR product from species coli is not cleaved by Dra I, the PCR product from species upsaliensis is only cleaved by Dde I and PCR product from lari is only cleaved by Alu I. It is a straightforward matter for a person skilled in the art to identify whether the PCR product is cleaved by one or more of the above endonucleases and thus the method enables simple speciation of campylobacter into jejuni, coli, upsaliensis or lari.

The embodiments of the first aspect of the invention described above form embodiments of the second aspect of the invention also, provided that primers are selected so as to be adapted to amplify at least nucleotides 124-196 of sequence ID NO:1, or a campylobacter sequence corresponding thereto.

In a preferred embodiment of the second aspect the primers consist of at least 12 contiguous nucleotides from sequence ID NO:s 4 and 5. Where the primers are sequences ID NO:s 4 and 5 the PCR product is 256bp and the respective products of cleavage by Alu I, Dra I and Dde I differentiate between jejuni, coli, upsaliensis and lari.

In a further embodiment of the invention, increased sensitivity and specificity for the detection of the presence of Campylobacter DNA, e.g. in food and liquid samples, is provided by the following additional methodologies:

1. A nested PCR has been developed, and is performed by an additional round of amplification using primer sequences international primer sequences ID4 and ID5. Two exemplary primer sequences are identified as Cru 0476 (SEQ ID NO:6) and Cru 0474 (SEQ ID NO:7). Following the second round of amplification, an amplicon of approximately 173pb is obtained in the presence of Campylobacter DNA. This DNA fragment

- 7 -

retains the sequences for the restriction endonucleases Alu 1, Dde 1, and Dra 1, thus still enabling the speciation of the contaminating campylobacter.

2. Additional increased sensitivity and specificity is optionally achieved by southern transfer of the amplified PCR products obtained using oligonucleotide primers ID4 and ID5, followed by hybridisation with an internal probe (e.g. SEQ ID NO:8 probe sequence). The probe sequence spans the restriction sites for speciation of the contaminating campylobacter and therefore restriction digest analysis can be used in conjunction with the probe hybridisation to confer additional specificity. The probe can be labelled, for example with digoxigenin, or radiolabelled.

The extraction procedures for food and environmental samples preferably use an internal standard to enable qualitative estimation of extraction efficiency and the effects of nonspecific inhibition. The PCR "MIMIC" (Clontech Laboratories, Palo Alto, California) is a form of competitive PCR in which a non-homologous neutral DNA fragment is engineered containing the same primer templates as the target DNA. The amplimer produced from this construct is a fragment either smaller or larger than the target product. Known amounts of construct are added to the PCR reaction, and compete for the same primers, acting as an internal standard. Where a mimic is used, the mimic sequence is capable of being amplified by the same primers that amplify, under PCR conditions, the campylobacter sequence. The mimic, if cleaved by restriction endonucleases, does not form fragments that interfere with detection and/or speciation of campylobacter-the mimic is said to be "neutral".

It is preferred to carry out the PCR steps of the invention also using a mimic. In an example, mimic DNA is added to the

- 8 -

sample and PCR is performed according to the invention. PCR product is analyzed. If mimic DNA ha been amplified, this indicates that the PCR reaction has occurred properly. product can then be tested for products that indicate presence of campylobacter. If no mimic DNA is amplified then this indicates PCR has not fully been carried out, or has been inhibited in some way.

It is further preferred to carry out PCR using mimic DNA of known and varying quantities. After amplification, the various results are compared and it is observed which of the results has comparable amounts of amplified mimic and (if present) campylobacter DNA. Thus, an estimate of the quantity of campylobacter DNA in the original sample is obtained.

The methods of the invention are further illustrated by the further embodiments of the invention described in the following Examples:-

Example 1

The PCR assay was developed by the following steps:

- Identification of a highly conserved, species specific 1> clone from a random library of Campylobacter jejuni insert fragments, cloned in the vector pBlueScript KS.
- Chain termination sequencing of the 1.9 kilobase fragment in both directions.
- Selection of presumptive primer pairs based on regions of equivalent G+C/A+T content, and low identity (prevention of 'primer-dimer').
- Optimisation of reaction parameters: Mg** concentration, 4> Taq enzyme source, buffer composition, annealing temperature,

- 9 -

cycling parameters.

Example 2

ASSESSMENT OF ASSAY SENSITIVITY AND SPECIFICITY

Using a single amplification (35 cycles, annealing temperature 50°C) we detected approx. 10 CFU/ml of Campylobacter jejuni.

We at this stringency, the assay was specific for Campylobacter jejuni, Campylobacter coli and Campylobacter upsaliensis. Using a lower annealing temperature (42°C), Campylobacter fetus and Campylobacter lari were also amplified.

The following procedures were used for PCR amplification of Campylobacter jejuni from milk and water samples.

- 1> cell lysis by boiling or freeze/thaw cycles, centrifuge, pcr supernatant directly.
- 2> Cell lysis by boiling, nucleic acid purification by phenol\chloroform extraction
- 3> cell lysis by guanidine isothiocyanate, nucleic acid purification using nuclease binding matrix ("isoquick").
- 4> Cell concentration using magnetic particles coated with anti-campylobacter igg, cell lysis by boiling.
- 5> concentration and immobilisation of cells on 0.2μm nitrocellulose filters ('solid-phase' pcr).
- 6> Cell concentration using affinity column purification
- 7> guanidium isothiocyanate nucleic acid extraction, with purification using silica bead matrix ('boom method')

- 11 -

EXTRACTION OF MILK SAMPLES

FOR PCR ANALYSIS

WARM MILK TO 37°C

CENTRIFUGE @ 3,000xg, 15 MINUTES

CHILL ON ICE. SEPARATE MILK AND CREAM

CREAM

MILK

EMULSIFY IN 10 VOLUMES OF WARM PBS

CENTRIFUGE @ 9,000xg, 15 MINUTES

DISCARD SUPERNATANT

DISCARD SUPERNATANT

RESUSPEND MILK AND CREAM PELLETS IN 5 VOLUMES OF PBS. POOL EXTRACTS.

BOIL FOR 10 MINUTES

CENTRIFUGE @ 14,000xg, 5 MINUTES

EXTRACT DNA WITH SILICA-BASED PURIFICATION MATRIX

ELUTE NUCLEIC ACIDS WITH 2x50µl PURE WATER

PCR NEAT SAMPLE, AND 10-FOLD SERIAL DILUTIONS

EXTRACTION OF WATER SAMPLES

PRE-FILTRATION THROUGH 30µm WHATMAN FILTER

CENTRIFUGE @ 9,000×g, 15 MINUTES

WASH PELLET x2, 1ML PBS

RESUSPEND IN 1ML STERILE WATER

BOIL, 10 MINUTES

EXTRACT DNA WITH SILICA-BASED

PURIFICATION MATRIX

Example 4

We observed the following differentiation of Campylobacter species using PCR primes SEQ ID NO:4 and SEQ ID NO:5 and restriction endonucleases Alu I, Dra I and Dde I.

	PCR product digested with						
Species	Alu I	Dra I	Dđe I				
C. jejuni	+	+	+				
C. coli	+	-	+				
C. upsaliensis	-	-	. +				
c. lari	+	-					

We further observed the following fragment sizes for different species.

Species		Restriction e	PCR amplimers				
		Alu I		Dde I	Dra I		
Thermophilic/ enteropathogenic		Fragment sizes (bp)		Fragment sizes (bp)		Fragment sizes (bp)	
C. jejuni	2	108, 148	2	83, 173	2	123, 133	
C. jejuni (hippurate + ve)	2	108, 148	2	83, 173	2	123, 133	
C. coli	2	108, 148	2	83, 173	1	256	
C. lari	2	108, 148	1	256	1	256	
C. upsaliensis	1	256	3	30,83,143	1	256	

The results are also illustrated in Fig. 3 where bonds were not visible by eye they were detected by use of radiolabels.

To test the specificity of campylobacter detection we used PCR primers on laboratory samples containing a wide range of organisms. The primers were SEQ ID NO:s 4 and 5, PCR product size in brackets:

	Annealing tem	perature of pr	
Species	37°C	42°C	50°C
C. jejuni	+ (256)	+	+
C. coli	+	+	+
C. upsaliensis	+	+	+
C. fetus	+	±	-
C. lari	+	±	-
C. mucosalis	±	-	-
C. sputorum	±	-	
Achromobacter sp.	-	<u>-</u> .	
Acinetobacter calcoac.	± (multiple)	-	-
Acinetobacter sp.	± (multiple)	_	
Aeromonas hydrophila		-	-
Citrobacter freundii	-	-	-
Enterobact. aerogenes	-	-	-
Enterobact. agglomerans	± (500)	-	-
Enterobacter cloacae	-	-	-
Escherichia coli	-		-
Flavobacterium	_	-	-

Species	37°C	42°C	50°C
Klebsiella aerogenes	A-	-	<u>-</u>
Klebsiella oxytoca	± (500)	-	-
Proteus mirabilis	-	_	_
Proteus morganii	_	_	_
Providencia stuartii	-	_	-
Pseudomonas aeroginosa	- sapr	-	_
Pseudomonas maltophilia	_		_
Pseudomonas pickettii	-	-	<u></u>
Salmonella enteritidis	-	- .	-
Salmonella typhimurium	· -	-	-
Serratia marcescens	-	-	-
Serratia liquefaciens	-	-	-
Shigella dysenteriae	·-	-	-
Shigella sonnei	-		· —
Vibrio cholera		-	-
Vibrio furnassii	± (1000)	-	_
Vibrio parahaemolyticus	± (180)	±	_
Yersinia enterocolitica	_	_	_
Oxford staphlococcus	± (300)	± -	_

Using standard culture techniques (published by Bolton F J, et al, J. Appl. Bacteriol., 1983, vol. 54, pages 115-125) we compared the detection of campylobacter jejuni by culture with detection by the method of the invention (using primers SEQ ID NO:s 4 and 5) against time.

The success of culture detection declined over the time of the comparison, no culturable organisms being found remaining in the sample after 26 days - thus at this point detection by culture indicated no campylobacter present.

By contrast, using the PCR method of the invention we were still able to detect campylobacter DNA in a sample 42 days old. The results are illustrated in Figure 1.

To confirm the accuracy of the PCR method of the invention we tested many samples that contained known species of campylobacter. The results, illustrated in figs. 4-11, confirm the method is completely accurate for all samples tested, and correctly identified each one by species.

FEATURES OF THE PCR ASSAY FOR Campylobacter jejuni

It allows rapid and sensitive detection of Campylobacter jejuni from environmental samples,

provides a semi-quantitative indication of the bacterial load, and determines whether samples are contaminated with Campylobacter jejuni, coli, upsaliensis or lari.

The method is of use for examining epidemiology of campylobacter infection such as a) seasonal peak, b) inverse correlation of surface water viable counts with human disease, c) role of water supply in (re)infection of broiler flocks with Campylobacter jejuni, d) contamination of foodstuffs at the point of sale, and e) determine origin of sporadic human infections.

Thus, a novel method incorporating polymerise chain reaction assay has been developed for the detection of campylobacter in clinical, environmental and food samples, such as milk and water samples. The assay is rapid, highly sensitive, and specific for Campylobacter sp. Simple restriction analysis of the PCR product allows speciation between Campylobacter jejuni, coli upsaliensis and lari.

Description of Drawings

- Fig. 1 shows a comparison of culturability of campylobacter jejuni against time with detection of campylobacter jejuni using PCR of the invention;
- Fig. 2 shows the sequence of open reading frame "C" from insert fragment pBSKSCJ19B with primer/nested primer locations, and restriction sites;
- Fig. 3 shows restriction enzyme analysis of PCR products amplified from C. jejuni, coli and upsaliensis.
- Fig. 4-11 shows the results of carrying out the PCR method of the invention on samples containing a wide ranges of known isolates. "P"=Penner Serotype Reference strains. "L"=Lior Serotype Reference Strains. Others are laboratory isolates. Standard size markers are on the gel ends.

Fig.	4	c.	Jejuni.	Alu	I	digest.	
Fig.	5	c.	Jejuni.	Dde	I	digest.	
Fig.	6	c.	Jejuni.			digest.	
Fig.	7	c.	upsaliensis.	Alu	I	and Dde I digest.	•
Fig.	8	c.	upsaliensis.			digest.	
Fig.	9	c.	Coli.	Alu	Į	digest	
Fig.	10	c.	Coli.	Dde	I	digest	
Fig.	11.	c.	Coli.	Dra	I	digest	

Sequence ID No. 1

- 1 accaacagcc attaaaaatc ttgactcagc catactcact ttaagaacac tggttgtcgg taatttttag aactgagtcg gtatgagtga aattcttgtg
- 51 gcggacctat ataataccgt tgcccaaatc cctgaaagca taaaaccaaa cgcctggata tattatggca acgggtttag ggactttcgt attttggttt
- 101 aatcacact gaagtatgaa gtggtctaag tcttgaaaaa gtggcatatt ttagtgtgga cttcatactt caccagattc agaacttttt caccgtataa
- 151 gtcctggtaa ataatttaaa ttaggatatg ccatttgaaa agctataaga caggaccatt tattaaattt aatcctatac ggtaaacttt tcgatattct
- 201 gttcctatag ccataccaac aatgccaaac aatatggtcg caaacataaa caaggatatc ggtatggttg ttacggtttg ttataccagc gtttgtattt
- 251 atatcttgca accgtatagt cgtaatttaa tacattacct ggatgcatcg tatagaacgt tggcatatca gcattaaatt atgtaatgga cctacgtagc
- 301 actttctcct taaaattttt gataacaaga gaagattata gaatattaat tgaaagagga attttaaaaa ctattgttct cttctaatat cttataatta
- 351 tatacatttt ttcttaaaaa tgat/aatttt gttaatcatt tgttatgttt atatgtaaaa aagaattttt acta ttaaaa caattagtaa acaatacaaa
- 401 tatattttaa ggctaaatca gtcttattta ttgatattta tcttataacc atataaatt ccgatttagt cagaataaat aactataaat agaatattgg
- 451 taaacttgtc acatttttta taaaatcttc acccacttta tctcttactc atttgaacag tgtaaaaaat attttagaag tgggtgaaat agagaatgag
- 501 tttttataaa agttcta{aca gcagtatcgc tcacatgt}c acctatccaa aaaaatattt tcaagat tgt cgtcatagcg agtgtaca g tggataggtt
- 551 acatttttct taatatcttc atgcaaaacc aaagctccag gttgctttaa tgtaaaaaga attatagaag tacgttttgg tttcgaggtc caacgaaatt
- 601 aagcaaagaa ataaaagcca attcttttt agttaaaaca atttctccac ttcgtttctt tattttcggt taagaaaaaa tcaattttgt taaagaggtg
- 651 cactgtaaat taaagttcgt ttatttttgt taaattgata ttcttcagaa gtgacattta atttcaagca aataaaaaca atttaactat aagaagtctt
- 701 atttttacaa gcatatttgc ttcaattttt tcacctatca gataatctaa taaaaatgtt cgtataaacg aagttaaaaa agtggatagt ctattagatt
- 751 aactttaaac aactcttcta tatcaacagg tttaatcaaa tatttatcta ttgaaatttg ttgagaagat atagttgtcc aaattagttt ataaatagat

Sequence ID No. 1 (cont)

- 801 taccaatatc aatagaacgc aaaagtctct ctt tctctga atacgcacta atggttatag ttatcttgcg ttttcagaga gaa{agagact tatgcgtgat
- 851 aga acaacaa ttgggacatc atctgaaatt tctttaatct ctcttgccat tct tgttgaa aaccctgtag tagactttaa agaaattaga gagaacggta
- 901 atccagtcca tccataatag gcatagcaat atctgtgata actaaatctg taggtcaggt aggtattatc cgtatcgtta tagacactat tgatttagac
- 951 gcttaaattt tttaaatttt ttaagcccct catctccatt ttgagctccg cgaatttaaa aaatttaaaa aattcggggt gtagaggtaa aactcgaggc
- 1001 attactttac taaagcgttc gcttaatata ttaatcattg attctctagc taatgaaatg atttcgcaag cgaattatat aattagtaac taagagatcg
- 1051 cttaacctca tcttcaacta ctaatattat taattcttta cattcttgtg gaattggagt agaagttgat gattataata attaagaaat gtaagaacac
- 1101 acat/ttctac tctaccctct cttttagttt taaaaatatc tcaaaacaag tgta aagatg agatgggaga gaaaatcaaa atttttatag agttttgttc
- 1151 ccccgtcttt tccattttta acttttattt ttccttggaa actttcgata ggggcagaaa aggtaaaaat tgaaaataaa aaggaacctt tgaaagctat
- 1201 atttgtctac ttatataaag tcctactcct ataccttgac taggatgttt taaacagatg aatatatttc aggatgagga tatggaactg atcctacaaa
- 1251 tgttgtaaaa taaggttgaa aaattttatc taaattttct ttatcaatcc acaacatttt attccaactt tttaaaatag atttaaaaga aatagttagg
- 1301 caccagcatt atcttttatt gtaattttca gataattttt tccaaatttt gtggtcgtaa tagaaaataa cattaaaagt ctattaaaaa aggtttaaaa
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- 1401 tgaatttaaa atcaaattaa gaaaaactct tattaaacca ttctcatatg acttaaattt tagtttaatt ctttttgaga ataatttggt aagagtatac
- 1451 ccaaaacttc ataatcactt ttcgaaacaa tattaatatt tacatgattt ggttttgaag tattagtgaa aagctttgtt ataattataa atgtactaaa
- 1501 ttttctatag tttcaaaagc aatttccaag gctttattta aagtctcttt aaagatatc aaagttttcg ttaaaggttc cgaaataaat ttcagagaaa
- 1551 tataaataca cactgctcta ctcctttgtt aaacaaagtt ctaaacacat
- 1601 caattgtttc tgacatattt ttaatcatat cttttgattg tgagtaaatt gttaacaaag actgtataaa aattagtata gaaaactaac actcatttaa
- 1651 tcagcaaatc ctttttcatc tttaagattt tgcttcattt gaaacatggc agacgtttag gaaaaagtag aaattctaaa acgaagtaaa ctttgtaccg

Sequence ID No. 1 (cont)

- 1701 aataccgagc tcatttaacg gttgtctcca ttgatgtgct atatcactaa ttatggctcg agtaaattgc caacagaggt aactacacga tatagtgatt
- 1751 tcatttgttc taatgaagat ttcaaaatct cttcatatgc tattttaata agtaaacaag attacttcta aagttttaga gaagtatacg ataaaattat
- 1801 tettttteat tttttteaa ggeaatttge attttttet caaatttttt agaaaaagta aaaaaaggtt eegttaaacg taaaaaaaga gtttaaaaaa
- 1851 acctaactgt ataaattctt gttggtgatt tttaactgta ttttcaagat tggattgaca tatttaagaa caaccactaa aaattgacat aaaagaacta
- 1901 taatacttaa ttctcttaat ttagcgtgat ttagagcaag ctcttcat attatgaatt aagagaatta aatcgcacta aatctcgttc gagaagta

Sequence ID No. 2

5' TCTTAGTGCG TATTCAGAGA 3'

Sequence ID No. 3

5' ACAGCAGTAT CGCTCACATG T 3'

Sequence ID No. 4

5' AGAACACGCG GACCTATATA 3' (also referred to as B04263)

Sequence ID No. 5

5' CGATGCATCC AGGTAATGTA T 3'
(also reffered to as B04264)

Primer Sequences

Internal

Cru 0476 (SEQ ID NO: 6)

5' atcacacctgaagtatga3' 18 mer

Cru 0474 (SEQ ID NO: 7)

5' tacgactatacggttgca3' 18 mer

Amplimer size: 173 base pairs

SEQ ID NO: 8

Probe sequence (173 bp amplimer from nested primers)

tacgactatacggttgcaagatattttatgtttgcg
----+
atgctgatatgccaacgttctataaaatacaaacgc

YDYTVARYFMFA

accatattgtttggcattgttggtatggct
----+1740
tggtataacaaaccgtaacaaccataccga
T I L F G I V G M A -

Alu I
ataggaactcttatagcttttcaaatggca
1741-----tatccttgagaatatcgaaaagtttaccgt
I G T L I A F Q M A -

Dra I
tatcctaatttaaattatttaccaggacaa
----+1800
ataggattaaatttaataaatggtcctgtt
Y P N L N Y L P G Q -

Dde I

tatgccactttttcaagacttagaccacttcatacttcaggtgtgat
1801----atacggtgaaaaagttctgaatctggtgaagtatgaagtccacacta

Y A T F S R L R P L H T S G V I

PCR MIMIC primers and sequence

Primer 1 Cru 0477 (SEQ ID NO: 9)

5' agaacacgcggacctatatacgcaagtgaaatctcctccg 3' 40 mer

Primer 2 Cru 0660 (SEQ ID NO: 10)

5' cgatgcatccaggtaatgtattctgtcaatgcagtttgtag 3' 41mer

MIMIC SEQUENCE (SEQ ID NO: 11)

agaacacgcg gacctatata cgcaagtgaa atctcctccg
tcttggagaa gggagagcgt ttgccccage taccattgat
gtgtacatga tcatggtcaa atgctggatg attgatgcag
acagccgtcc caagtttcgt gagctgattg cagagttctc
caaaatggct cgtgaccctc cccgctatct tgttatacag
ggagatgaaa ggatgcactt gcctagccct acagattcca
agttttatcg caccctgatg gaggaggagg acatggaaga
cattgtggat gcagatgagt atcttgtccc acaccagggc
tttttcaaca tgccctctac atctcggact cctcttctga
gttcattgag cgctactagc aacaattctg ctacaaactg
cattgacaga 3'

CLAIMS:

1. A method of testing for the presence of campylobacter in a clinical, environmental or food sample comprising the steps of:-

performing polymerise chain reaction (PCR) using primers adapted to amplify a region selected from (a) a sequence of at least 72 base pairs from sequence ID No. 1 and (b) a sequence having sufficient homology with (a) such that formation of PCR product is correlated with presence of campylobacter; and determining if any PCR product is formed.

- 2. A method according to claim 1 in which sequence (b) has at least 75% homology with (a), preferably at least 90% homology and more preferably at least 95% homology.
- 3. A method according to Claim 1 or 2 in which the primers comprise at least 12 nucleotides.
- 4. A method according to Claim 3 in which the primers are 19-22 nucleotides in length.
- 5. A method according to Claim 3 or Claim 4 in which the primers consist of at least 12 nucleotides selected from:
 - (1) sequence ID No.2 and sequence ID No.3,
 - (2) sequence ID No.4 and sequence ID No.5, and
 - (3) sequences having sufficient homology with (1) or (2) such that formation of PCR product is correlated with presence of campylobacter.

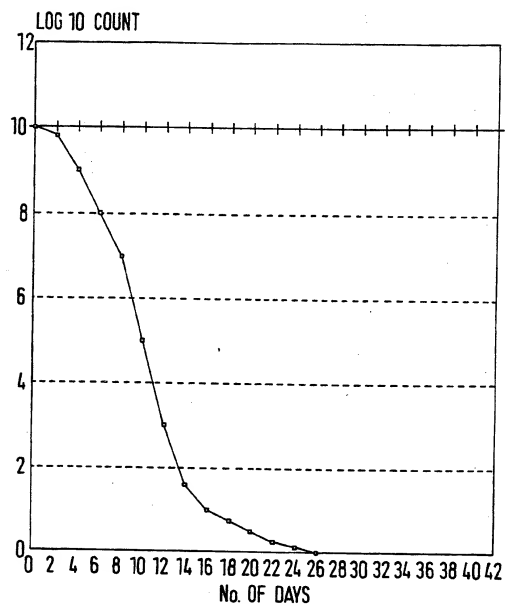
- 6. A method according to any previous claim in which the temperature of the PCR is sufficiently high to prevent the primers annealing with non-campylobacter DNA.
- 7. A method according to Claim 6 in which the PCR temperature is at least 40°C .
- 8. A method according to Claim 7 in which the PCR temperature is at least 45°C .
- 9. A method according to Claim 8 in which the PCR temperature is $48-52^{\circ}\text{C}$.
- 10. A method of distinguishing between campylobacter species jejuni, coli, upsaliensis and lari in a DNA containing sample by performing PCR utilizing primers capable of amplifying a selected campylobacter DNA sequence, said sequence having restriction endonuclease sites specifically associated with the different campylobacter species, obtaining PCR product and then testing for digestion of the PCR product by the specific restriction endonucleases.
- 11. A method according to claim 10 wherein the selected campylobacter sequence is differentially cleaved by the restriction endonucleases Alu I, Dra I and Dde I
- 12. A method of distinguishing between campylobacter species jejuni, coli and upsaliensis comprising the steps of:-

performing polymerise chain reaction (PCR) using primers adapted to amplify a region of DNA sequence ID No.1 that includes nucleotides 124-196, or using primers adapted to amplify a region of campylobacter DNA corresponding thereto; and

testing the PCR product for digestion by restriction endonucleases Alu I, Dra I and Dde I.

- 13. A method according to Claim 10, 11 or 12 in which campylobacter jejuni is characterised by cleavage by all three endonucleases, coli is characterised by loss of cleavage by Dra I endonucleases, upsaliensis is characterised by loss of cleavage by Alu I and Dra I endonucleases and lari is characterised by cleavage by Alu I only.
- 14. A method according to any of Claims 10 13 characterised by the features of any of claims 3-9.
- 15. Use of restriction endonucleases Alu I, Dra I and Dde I in differentiating between campylobacter species jejuni, coli upsaliensis and lari.
- 16. A PCR primer having the sequence of sequence ID No. 2.
- 17. A PCR primer having the sequence of sequence ID No. 3.
- 18. A PCR primer having the sequence of sequence ID No. 4.
- 19. A PCR primer having the sequence of sequence ID No. 5.
- 20. A PCR primer having the sequence of sequence ID No. 6, 7, 9 or 10.
- 21. A kit for detecting campylobacter comprising one or more reagents for carrying out the method of any of claims 1-9.
- 22. A kit for determining campylobacter species comprising one or more reagents for carrying out the method of any of Claims 10-15.

1/8
CULTURABILITY AND PCR DETECTION OF
C. JEJUNI IN A POND WATER MICROCOSM



- SURFACE VIABLE COUNT + DILN DETECTED BY PCR

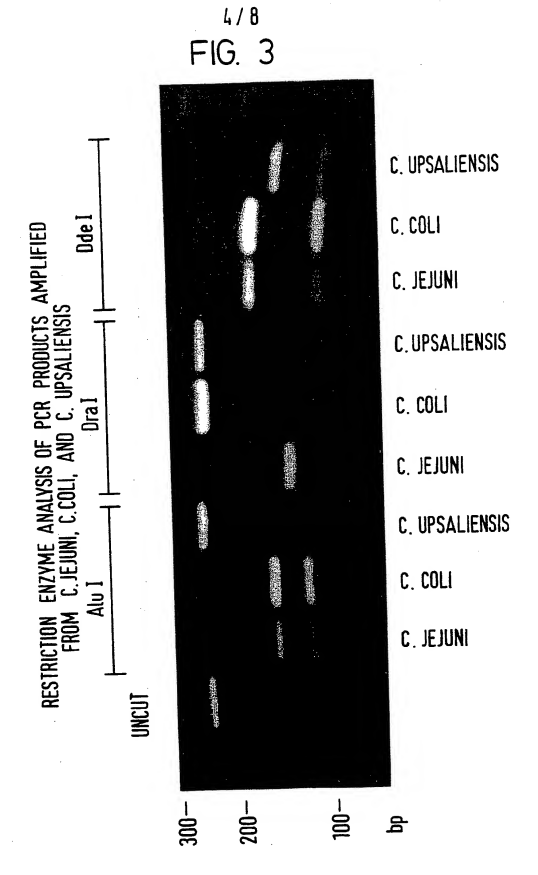
FIG. 1

			FIG. 2(I)		2/8						
			(1)		· .						
		1	•		1			1			,
tgattaacaaaattatcatttttaagaaaaatgtataattaat	attottttttacatattaattataagatattagaagag	i,	gaaagtcgatgcaggtaatgtattaaattacgac	aacaatagtttttaaaattcctctttcagctacgtaggtccattacataatttaatgctg	Q	acggttgcaagatattttatgtttgcgaccatattgtttggcattgttggtatggct	cga	Æ	Caa	gtt	a
ctt	gaa	H	tac !	atg	×	atg	tac	Σ	gga	act	0
aat	tta	z	aat	tta	z	ggt	CCa	_ ა	Dra I aaatggcatatcctaa tttaaa ttatttaccaggacaa	tttaccgtataggatt aaatt taataaatggtcctgtt	d
tctata	ata.	≻	t a	aat	Н	ttgttgg t	Zaa(>	t tac	aat	
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tat	ate		lato	tac	>	tattgtttggca	cgt	H	H Hatt	taa	>
tae	att	* _	gta	cat	Z	ttg	aac	Ö	Dra :ttaa	att	Z
ataatta	tta	z	Ca3	gto	IJ	tgt	aca		att	taa	1
tat	ata	* ≯.	8 1 0 1	tag	₽4	tat	ata	-	i ta	gat	2
atg	tac	Σ_	tge	acg	Ħ	CCa	ggt	H	a tc	tag	ָם i
333	ר רר	× z	- t	jct	Σ	ttgcgacca	jcti	H	; at	tai	>
Jaa	it t	ы _ ж	ag të	ည်သင်	S	tgc	acç	A	ggc	SSS	A
aac	tto	æ	Jaac	itt	×	gtt	caa	<u>le</u>	aat	tta	Σ
tt	laae	* 17	gag	ctc	E	tat	ata	Σ	1 C	agt	a
att	taa	Er Er	aag	tto	×	gatatttta	aaa	[24	+ t t	a aa	De,
atc	tag	π S	tt	aaa	H	ata	tat	×	Alu I agctt	tcg	A
att	taa	r K	aat	tta	н	aag	ttc	×	ו ניל	ata	
988	t t	Z	3883 -+-	tt	×	9	acg1	Æ	; ; ;	agad	TLI
taacaaattatcattt	ttg	Z Z Z	atc	ag	Ø	ggttgcaa	Cas	>	lact	tge	E
tt	actaattgttttaatagtaaa	*	ttgttatcaaaaattttaaggaga	aat	1	308	atatgccaacgttctataaaatacaaacgctggtataacaaaccgtaacaaccataccga	E	Alu I ataggaactettatagetttteaaatggeatateetaatttaaaattaeeaggaeaa	tatccttgagaata tcga aaag	6
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tatgccactttttcaagactacttcatacttcaggtgtgatttttggttttatg	atacggtgaaaaagttct gaatc tggtga agtatgaagtccac taaaaaccaaaatac	YATFSRLRPLHTSGVIFGFM -	ctttcagggatttgggcaacggtattatataggtccgcgtgttcttaaagtgagtatggc	gaaagtccctaaaacccgttgccataatatccaggcgcacaagaatttcactcataccg	LSGIWATVLYRSACS*SEYG -	tgagtcaagatttttaatggctgttggt 	
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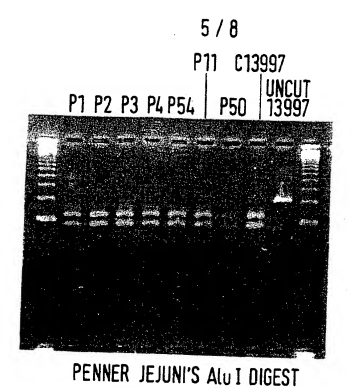


FIG. 4 SIZE MARKERS ON ENDS

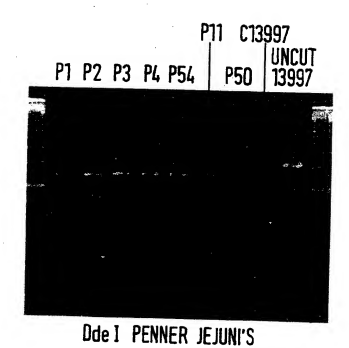


FIG. 5

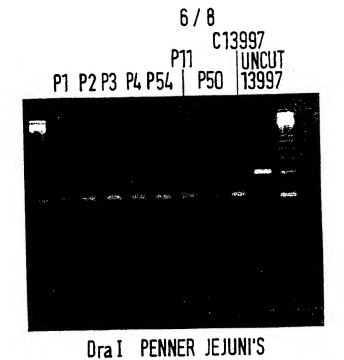
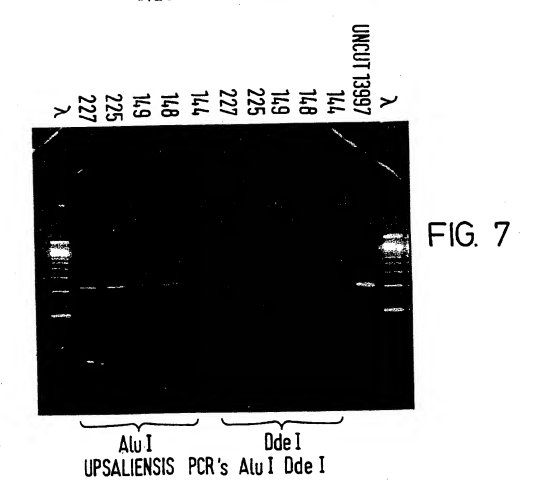


FIG. 6



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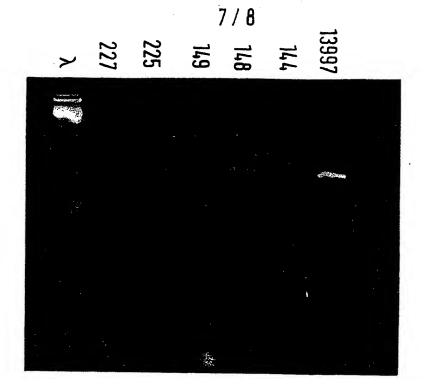


FIG. 8

UPSALIENSIS PCR's Dra I CUT

P5 P24 P25 P59 L8 L2 COLI

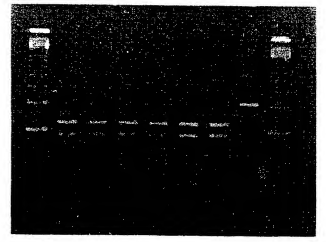


FIG. 9

Alu I - PENNER / LIOR COLI'S

8/8 P5 P24 P25 P59 L8 L2 COLI

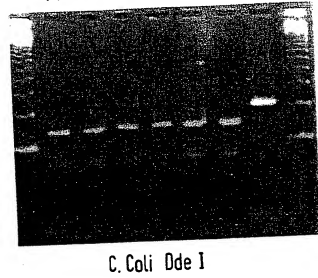
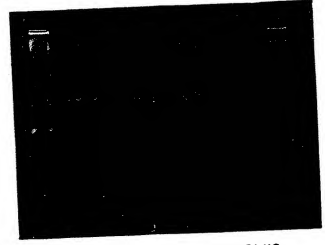


FIG. 10

FIG. 11

P5 P24 P25 P59 L8 L21 COLI



Dra I PENNER / LIOR COLI'S

INTERNATIONAL SEARCH REPORT

International application No. PCT/GR 94/01967

			PCT/GB 94	I/01967
IPC 6	SSIFICATION OF SUBJECT MATTER C12Q1/68			
According	g to International Patent Classification (IPC) or to both nationa	classification and IPC		
B. FIELD	DS SEARCHED			
IPC 6	documentation searched (classification system followed by cla $C12Q$	ssification symbols)		· · ·
Document	ation searched other than minimum documentation to the exten	t that such documents are include	ed in the fields se	arched
Electronic	data base consulted during the international search (name of da	ita base and, where practical, sea	rch terms used)	
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of	the colours access		
	appropriate, u	die reievant passages		Relevant to claim No.
X	J CLIN MICROBIOL 31 (6). 1993. NACHAMKIN I et al 'FLAGELLIN G OF CAMPYLOBACTER -JEJUNI BY ST FRAGMENT LENGTH POLYMORPHISM A see the whole document	ENE TYPING RICTION		10-15
X	EP,A,O 350 392 (IRE-MEDGENIX S January 1990 see claims 12,13; example 5			15
A	LETTERS IN APPLIED MICROBIOLOG 1993. 235-237. Birkenhead D et al 'PCR for the and typing of campylobacters.'			10-15
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X Further	er documents are listed in the continuation of box C.	X Patent family memb	ers are listed in a	nnex.
A* document consider earlier do filing da L' document which is citation of document other me document ager that	at which may throw doubts on priority claim(s) or a cited to establish the publication date of another or other special reason (as specified) at referring to an oral disclosure, use, exhibition or eans at published prior to the international filing date but in the priority date claimed challenged completion of the international search	"T' later document published or priority date and not cited to understand the pinvention "X' document of particular recannot be considered no involve an inventive step "Y' document of particular recannot be considered to document is combined we ments, such combination in the art. '&' document member of the	I after the internation conflict with the minciple or theory cleavance; the claim vel or cannot be a when the documelet an involve an invention one or more their obvious to same patent famernational search	ational filing date the application but y underlying the timed invention considered to the time to taken alone med invention tive step when the other such docu- to a person skilled
	January 1995 illing address of the ISA Firmness Peters Office P.P. Cale P.A.	10.02.9)5 	·
	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijawijk Tcl. (+31-70) 340-2040, Tx. 31 651 epo nl, Fac: (+31-70) 340-3016	Molina Gala	an,-E	

INTERNATIONAL SEARCH REPORT

International application No. PCT/GB 94/01967

		PCT/GB 94/01967
(Continuati	(ca) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.
ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Vetesant & comm 1.00
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A	APPLIED AND ENVIRONMENTAL MICROBIOLOGY, vol.58, no.12, December 1992 pages 3804 - 3808 GIESENDORF ET AL. 'Rapid and sensitive detection of C. spp. in chicken products by using PCR' cited in the application	

INTERNATIONAL SEARCH REPORT

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International application No. PCT/GB 94/01967

	Determs do				34701307		
	Patent document cited in search report	Publication date	Patent memb		Publication date		
	EP-A-0350392	10-01-90	FR-A- JP-A-	2633941	12-01-90		
1		***		2154700	14-06-90		

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